## PATENT COOPERATION TREATY

To:

PCT

NOTICE INFORMING THE APPLICANT OF THE

COMMUNICATION OF THE INTERNATIONAL

APPLICATION TO THE DESIGNATED OFFICES

(PCT Rule 47.1(c), first sentence)

From the INTERNATIONAL BUREAU

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Birch

Date of mailing (day/month/year) 29 January 1998 (29.01.98)

Applicant's or agent's file reference

97040PCT

IMPORTANT NOTICE

International application No. PCT/JP97/02484

International filing date (day/month/year)
17 July 1997 (17.07.97)

Priority date (day/month/year) 18 July 1996 (18.07.96)

**Applicant** 

KAO CORPORATION et al

 Notice is hereby given that the International Bureau has communicated, as provided in Article 20, the international application to the following designated Offices on the date indicated above as the date of mailing of this Notice:

CA, EP, JP, US

In accordance with Rule 47.1(c), third sentence, those Offices will accept the present Notice as conclusive evidence that the communication of the international application has duly taken place on the date of mailing indicated above and no copy of the international application is required to be furnished by the applicant to the designated Office(s).

2. The following designated Offices have waived the requirement for such a communication at this time:

None

The communication will be made to those Offices only upon their request. Furthermore, those Offices do not require the applicant to furnish a copy of the international application (Rule 49.1(a-bis)).

3. Enclosed with this Notice is a copy of the international application as published by the International Bureau on 29 January 1998 (29.01.98) under No. WO 98/03730

#### REMINDER REGARDING CHAPTER II (Article 31(2)(a) and Rule 54.2)

If the applicant wishes to postpone entry into the national phase until 30 months (or later in some Offices) from the priority date, a demand for international preliminary examination must be filed with the competent International Preliminary Examining Authority before the expiration of 19 months from the priority date.

It is the applicant's sole responsibility to monitor the 19-month time limit.

Note that only an applicant who is a national or resident of a PCT Contracting State which is bound by Chapter II has the right to file a demand for international preliminary examination.

## REMINDER REGARDING ENTRY INTO THE NATIONAL PHASE (Article 22 or 39(1))

If the applicant wishes to proceed with the international application in the national phase, he must, within 20 months or 30 months, or later in some Offices, perform the acts referred to therein before each designated or elected Office.

For further important information on the time limits and acts to be performed for entering the national phase, see the Annex to Form PCT/IB/301 (Notification of Receipt of Record Copy) and Volume II of the PCT Applicant's Guide.

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland Authorized officer

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Form PCT/IB/308 (July 1996)

## PATENT COOPERATION TREATY

### **PCT**

## INFORMATION CONCERNING ELECTED OFFICES NOTIFIED OF THEIR ELECTION

(PCT Rule 61.3)

#### From the INTERNATIONAL BUREAU

To

FURUYA, Kaoru Nihonbashi TM Building 1-8-11, Nihonbashi-Horidomecho Chuo-ku Tokyo 103 JAPON

Date of mailing (day/month/year)

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**Applicant** 

KAO CORPORATION

1. The applicant is hereby informed that the International Bureau has, according to Article 31(7), notified each of the following Offices of its election:

EP:AT,BE,CH,DE,DK,ES,FI,FR,GB,GR,IE,IT,LU,MC,NL,PT,SE National:CA,JP,US

2. The following Offices have waived the requirement for the notification of their election; the notification will be sent to them by the International Bureau only upon their request:

None

3. The applicant is reminded that he must enter the "national phase" before the expiration of 30 months from the priority date before each of the Offices listed above. This must be done by paying the national fee(s) and furnishing, if prescribed, a translation of the international application (Article 39(1)(a)), as well as, where applicable, by furnishing a translation of any annexes of the international preliminary examination report (Article 36(3)(b) and Rule 74.1).

Some offices have fixed time limits expiring later than the above-mentioned time limit. For detailed information about the applicable time limits and the acts to be performed upon entry into the national phase before a particular Office, see Volume II of the PCT Applicant's Guide.

The entry into the European regional phase is postponed until 31 months from the priority date for all States designated for the purposes of obtaining a European patent including, where applicable. ES which cannot be elected since it is not bound by Chapter II.

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

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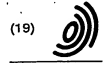
(54)Title: BULKING AGENTS FOR PAPER, HIGH-BULKINESS PULP SHEETS, AND PROCESS FOR THE PRODUCTION OF THE SHEETS

OF THE SHEETS

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(57) Abstract

Bulking agents for paper enabling the production of bulky pulp sheets without detriment to the strength of paper, which comprise impounds of general formula (1)  $RO(EO)_m (PO)_n H$  wherein R is  $C_6 - C_{22}$  linear or branched alkyl or alkenyl, or alkylaryl wherein the alkyl has 4 to 20 carbon atoms; E is ethylene; P is propylene; m and n each refer to an average number of molecules added and are a number of 0 to 20 and a number of 0 to 50, respectively; and the EO and PO units may be arranged in block or at random and either of EO and PO may be attached directly to RO.



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# (54) BULKING AGENTS FOR PAPER, HIGH-BULKINESS PULP SHEETS, AND PROCESS FOR THE PRODUCTION OF THE SHEETS

(57) A paper bulking promoter containing a compound represented by the following formula (1) with which a highly bulky sheet can be obtained without impairing paper strength:

$$RO(EO)_m(PO)_nH$$
 (1)

wherein R represents a linear or branched alkyl or alkenyl group having 6 to 22 carbon atoms or an alkylaryl group in which the alkyl group has 4 to 20 carbon atoms; E represents an ethylene group; P represents a propylene group; m indicates the average number of moles added in the range of 0≤m≤20; and n is a number in the range of 0≤n≤50; provided that the EO and PO groups may have any of block and random arrangements and may begin with any of EO and PO.

#### Description

#### Background of the Invention

#### Field of the Invention

[0001] This invention relates to a paper bulking promoter with which sheets of paper obtained from a pulp feedstock can be bulky without impairing paper strength.

#### Statement of Related Art

[0002] Recently, there is a desire for high-quality paper, e.g., paper excellent in printability and voluminousness. Since the printability and voluminousness of paper are closely related to the bulkiness thereof, various attempts have been made to improve bulkiness. Examples of such attempts include a method in which a crosslinked pulp is used (JP-A 4-185792, etc.) and a method in which a mixture of pulp with synthetic fibers is used as a feedstock for papermaking (JP-A 3-269199, etc.). Examples thereof further include a method in which spaces among pulp fibers are filled with a filler such as an inorganic (JP-A 3-124895, etc.) and a method in which spaces are formed (JP-A 5-230798, etc.). On the other hand, with respect to mechanical improvements, there is a report on an improvement in calendering, which comprises conducting calendering under milder conditions (JP-A 4-370298).

[0003] However, the use of a crosslinked pulp, synthetic fibers, etc. makes pulp recycling impossible, while the technique of merely filling pulp fiber spaces with a filler and the technique of forming spaces result in a considerable decrease in paper strength. Furthermore, the improvement in mechanical treatment produces only a limited effect and no satisfactory product has been obtained so far.

[0004] Also known is a method in which a bulking promoter is added during papermaking to impart bulkiness to the paper. Although fatty acid polyamide polyamines for use as such bulking promoters are on the market, use of these compounds results in a decrease in paper strength and no satisfactory performance has been obtained therewith.

#### Disclosure of the Invention

#### 30 Summary of the Invention

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[0005] The inventors have made intensive investigations in view of the problems described above. As a result, they have found that by incorporating a specific alcohol and/or a polyoxyalkylene adduct thereof into a pulp feedstock, e.g., a pulp slurry, in the papermaking step, the sheet made from the feedstock can have low density (improved bulkiness) without detriment to paper strength. This invention has thus been achieved.

[0006] Namely, this invention provides a paper bulking promoter which comprises a compound represented by the following formula

#### RO(XO)<sub>a</sub>H

wherein R represents a linear or branched alkyl or alkenyl group having 6 to 22 carbon atoms or an alkylaryl group in which the alkyl group has 4 to 20 carbon atoms; X's are the same or different and each represents a linear or branched alkylene group having 2 or 3 carbon atoms; and a indicates the average number of moles added in the range of 0≤a≤20. [0007] This invention relates to a paper bulking promoter containing at least one compound (1) represented by formula (1):

$$RO(EO)_{m}(PO)_{n}H$$
 (1)

wherein R represents a linear or branched alkyl or alkenyl group having 6 to 22 carbon atoms or an alkylaryl group in which the alkyl group has 4 to 20 carbon atoms; E represents an ethylene group; P represents a propylene group; and m and n respectively indicate the average number of moles added in the ranges of  $0 \le m \le 20$  and  $0 \le n \le 50$ ; provided that  $(EO)_m(PO)_n$  may have any of block and random arrangements and may begin with any of EO and PO.

[0008] In formula (1), R is preferably a linear or branched alkyl or alkenyl group having 8 to 18 carbon atoms.

[0009] The compound (1) includes one obtained by causing a mixture of two or more ROH's to add at least either of EO and PO. An alcohol represented by ROH is also included.

[0010] This invention further provides a paper bulking promoter composition which comprises the compound (1) and at least one nonionic surfactant based on a polyhydric alcohol.

[0011] The nonionic surfactant based on a polyhydric alcohol is preferably at least one member selected among sugar

alcohol/EO (ethylene oxide, the same applies hereinafter) adducts, fatty acid esters of the adducts, fatty acid esters of sugar alcohols, sugar/EO adducts, fatty acid esters of the adducts, sugar/fatty acid esters, and fat/EO adducts. Especially preferred are fatty acid esters of sugar alcohol/EO adducts and fat/EO adducts.

[0012] The proportion of the compound (1) to the nonionic surfactant based on a polyhydric alcohol is desirably from 5/5 to 10/0 (by weight), preferably from 5/5 to 99/1 (by weight).

[0013] This invention also provides a process for producing a highly bulky pulp sheet which comprises adding the bulking promoter or composition in any of the papermaking process steps to thereby produce a highly bulky pulp sheet which has a bulk density lower by at least 5% than the product not containing the paper bulking promoter and a tearing strength as measured according to JIS P 8116 of at least 90% of that of the product. The highly bulky pulp sheet obtained by this process is also included. It is preferred that the bulking promoter or composition be mixed with water and a pulp feedstock to obtain a homogeneous slurry, which is subjected to sheet forming.

[0014] Further, an aqueous emulsion containing 10 to 100 wt.% of the bulking promoter or composition may be mixed with a pulp feedstock to obtain a homogeneous slurry, which is subjected to sheet forming.

[0015] In the bulking promoter composition, m and n are desirably in the ranges of 0≤m≤2 and 0≤n≤50, and the proportion of the compound (1) to the nonionic surfactant based on a polyhydric alcohol is desirably from 5/5 to 99/1 (by weight). Preferably, m and n each is zero and the proportion of the compound (1) to the nonionic surfactant based on a polyhydric alcohol is from 5/5 to 99/1 (by weight). The especially preferred range is from 7/3 to 95/5 (by weight).

[0016] This invention furthermore provides a highly bulky sheet which comprises the bulking promoter or composition and a pulp paper and which has a bulk density lower by at least 5% than the product not containing the paper bulking promoter and a tearing strength as measured according to JIS P 8116 of at least 90% of that of the product.

#### **Detailed Explanation of the Invention**

[0017] The term "paper bulking promoter" used herein means an agent with which a sheet of paper obtained from a pulp feedstock can have a larger thickness (can be bulkier) than that having the same basis weight obtained from the same amount of a pulp feedstock.

[0018] The compound represented by the formula (1) is one obtained by causing a higher alcohol, an alkylphenol, or the like in which the alkyl has 6 to 22 carbon atoms to add an alkylene oxide such as ethylene oxide (EO) or propylene oxide (PO). In this invention is used the compound in which the average number of moles of ethylene oxide added is in the range of 0≤m≤20. The range of the average number of moles added, m, is desirably 0≤m≤10, preferably 0≤m≤5. If m exceeds 20, the effect of imparting bulkiness to paper is lessened. Further, the compound used is one in which the average number of moles of propylene oxide (PO) added, n, is in the range of 0≤n≤50, preferably 0≤n≤20. When n exceeds 50, such a compound is economically disadvantageous although the decrease in performance is little.

[0019] R in the formula (1), which represents a linear or branched alkyl or alkenyl group having 6 to 22 carbon atoms or an alkylaryl group in which the alkyl has 4 to 20 carbon atoms, is preferably a linear or branched alkyl or alkenyl group having 8 to 18 carbon atoms. If R is an alkyl or alkenyl group in which the number of carbon atoms is outside the range of from 6 to 22 or if R is an alkylaryl group in which the number of carbon atoms of the alkyl group is outside the range of from 4 to 20, then the compound is less effective in imparting bulkiness to paper.

[0020] E and P in the formula (1) each represents a linear or branched alkylene group having 2 or 3 carbon atoms. Examples thereof include ethylene and propylene. When the group  $(EO)_m(PO)_n$  in the general formula (1) is composed of a combination of polyoxyethylene and polyoxypropylene, the  $C_2H_4O$  and  $C_3H_6O$  groups may be in any of random and block arrangements. In this case, the polyoxypropylene  $(C_3H_6O)$  group(s) account for preferably at least 50 mol%, especially preferably at least 70 mol%, of all groups added on the average. The alkylene oxide group bonded to R may begin with any of EO and PO.

[0021] The paper bulking promoter of this invention preferably further contains a nonionic surfactant based on a polyhydric alcohol. By the use of a combination of the compound represented by the formula (1) and a nonionic surfactant based on a polyhydric alcohol, the effect of this invention can be improved. In particular, in the case where the compound represented by the formula (1) is less apt to dissolve in water when used alone and is difficult to evenly mix with a pulp feedstock, e.g., a pulp or pulp slurry, for example, in the case where the number of moles of EO added is 2 or smaller, in particular 0, then the effect of the combined use of the two ingredients is enhanced when this compound represented by the formula (1) is emulsified with the nonionic surfactant based on a polyhydric alcohol, although the compound can be dispersed mechanically.

[0022] The nonionic surfactant based on a polyhydric alcohol is desirably at least one member selected among sugar alcohol/EO adducts or fatty acid esters thereof, fatty acid esters of sugar alcohols, sugar/EO adducts or fatty acid esters thereof, sugar/fatty acid esters, and fat/EO adducts. Preferably, the nonionic surfactant based on a polyhydric alcohol is either of a fatty acid ester of a sugar alcohol/EO adduct and a fat/EO adduct. Especially preferred is a combination of these.

#### · (1) Nonionic surfactants based on sugar alcohol

[0023] Examples of the nonionic surfactants based on a sugar alcohol include sugar alcohol/EO adducts, fatty acid esters of sugar alcohol/EO adducts, and fatty acid esters of sugar alcohols. The sugar alcohol as a component of a nonionic surfactant based on a polyhydric alcohol is an alcohol obtained from a monosaccharide such as a triose, tetrose, pentose, or hexose through reduction of the aldehyde or ketone group. Examples thereof include the glycerol derived from trioses, the erythritol and threitol derived from tetroses, the arabitol, ribitol, and xylitol derived from pentoses, and the sorbitol, mannitol, altrose, and galactitol derived from hexoses. The sugar alcohol/EO adducts are nonionic surfactants of the ether type. These adducts preferably are ether ester type nonionic surfactants derived from a sugar alcohol. In this case, part of the hydroxy groups of the sugar alcohol form an ester with a fatty acid. The fatty acid as a component of the fatty acid ester in a sugar alcohol/EO adduct may be any of saturated and unsaturated fatty acids each having 1 to 24, preferably 12 to 18, carbon atoms. Preferred is oleic acid. With respect to the degree of esterification of the sugar alcohol, the number of -OH groups which have undergone esterification may be any of from zero to all of the -OH groups, that is, the sugar alcohol may, for example, be in the form of a mono-, sesqui-, di-, or tri ester. However, the degree of esterification is preferably 1 to 3. In the sugar alcohol/EO adduct or its fatty acid ester, the average number of moles of EO added is 0 to 100, preferably 10 to 50. When the average number of moles of EO added is 0, this compound is a sugar alcohol/fatty acid ester, which type of nonionic surfactant may be used in this invention. Preferred examples of the nonionic surfactant based on a sugar alcohol for use in this invention are fatty acid esters of sugar alcohol/EO adducts. Most desirable among these are polyoxyethylene sorbitan/fatty acid esters.

## (2) Nonionic surfactants based on sugar

[0024] Examples of the nonionic surfactants based on a sugar include sugar/EO adducts, fatty acid esters of sugar/EO adducts, and sugar/fatty acid esters. The sugar may be a polysaccharide such as sucrose, besides any of the monosaccharides mentioned above with regard to the sugar alcohol. In the sugar/EO adducts also, the average number of moles of EO added is 0 to 100, preferably 10 to 50. When the average number of moles of EO added is 0, this compound is a sugar/fatty acid ester. Examples of the sugar/fatty acid ester include sucrose/fatty acid esters. Examples of the fatty acid as a component of the ester may be the same as those mentioned above.

#### (3) Fat/EO adducts

[0025] Examples of fats usable as starting materials for the fat/EO adducts include vegetable oils such as castor oil, coconut oil, palm oil, olive oil, soybean oil, rapeseed oil, and linseed oil, animal fats such as porcine fat and beef tallow, fish oils, hardened and semihardened oils obtained therefrom, and recovery oils obtained during the purification of these fats. The most desirable among these fats is hardened castor oil. In the fat/EO adducts, the average number of moles of EO added is 5 to 100, preferably 10 to 50.

[0026] When the compound represented by formula (1) is used in combination with the nonionic surfactant based on a polyhydric alcohol as described above, the proportion of the compound represented by formula (1) to the nonionic surfactant based on a polyhydric alcohol is from 5/5 to 99/1, preferably from 7/3 to 95/5 (by weight). In the case where the nonionic surfactant based on a polyhydric alcohol is also used, the compound represented by formula (1) and the nonionic surfactant based on a polyhydric alcohol may be used in the form of an emulsion or mixture prepared by adding these ingredients to water under agitation. For industrial use, the emulsion or mixture has a concentration of about 10 to 100% from the standpoint of profitability.

[0027] The bulking promoter of this invention is applicable to a variety of ordinary pulp feedstocks ranging from virgin pulps such as mechanical pulps and chemical pulps to pulps prepared from various waste papers. The point where the bulking promoter of this invention is added is not particularly limited as long as it is within the papermaking process steps. In a factory, for example, the bulking promoter is desirably added at a point where it can be evenly blended with a pulp feedstock, such as, the refiner, machine chest, or headbox. After the bulking promoter of this invention is added to a pulp feedstock, the resultant mixture is subjected as it is to sheet forming. The bulking promoter remains in the paper. The paper bulking promoter of this invention is added in an amount of 0.01 to 10 wt.%, preferably 0.1 to 5 wt.%, based on the pulp.

[0028] The pulp sheet obtained by using the paper bulking promoter of this invention has a bulk density (the measurement method is shown in the Examples given later) lower by desirably at least 5%, preferably at least 7% than the product not containing the paper bulking promoter and has a tearing strength as measured according to JIS P 8116 of desirably at least 90%, preferably at least 95% of that of the product.

#### Effect of the Invention

[0029] By adding the bulking promoter of this invention in papermaking, a highly bulky sheet can be obtained without impairing paper strength.

**Examples** 

[0030] This invention will be explained below in more detail by reference to Examples, but the invention should not be construed as being limited thereto. In the Examples, all parts and percents are based on weight unless otherwise indicated.

Examples 1 to 20 and Comparative Examples 1 to 6

[Pulp Feedstock]

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[0031] The deinked pulp and virgin pulp shown below were used as pulp feedstocks.

( Deinked pulp )

20 [0032] A deinked pulp was obtained in the following manner. Feedstock waste papers collected in the city (newspaper/leaflet: 70/30%) were cut into a size of 4 cm by 4 cm, and a given amount of the cut paper was introduced into a bench disintegrator. Thereto were added warm water, 1% (based on the feedstock) of sodium hydroxide, 3% (based on the feedstock) of sodium silicate, 3% (based on the feedstock) of a 30% aqueous hydrogen peroxide solution, and 0.3% (based on the feedstock) of EO/PO block adduct of beef tallow/glycerol (1:1), as a deinking agent, in which the amounts of EO and PO were respectively 70 and 10 (average number of moles added). The feedstock was disintegrated at 40°C for 10 minutes at a pulp concentration of 5%. The pulp slurry obtained was aged at 40°C for 60 minutes and then diluted with warm water to a pulp concentration of 1%. The diluted slurry was subjected to flotation at 40°C for 10 minutes. After the flotation, the slurry was washed with water and regulated to a concentration of 1% to prepare a deinked pulp (DIP) slurry. The DIP had a freeness of 220 ml.

(Virgin pulp)

[0033] A virgin pulp was prepared by cutting an LBKP (bleached hardwood pulp) into a size of 5 cm by 5 cm and disintegrating and beating a given amount of the cut LBKP with a beater at room temperature to give a 1% LBKP slurry. This LBKP had a freeness of 420 ml.

[Papermaking Method]

[0034] Each of the above 1% pulp slurries was weighed out in such an amount as to result in a sheet of paper having a basis weight of 60 g/m². The pH thereof was adjusted to 4.5 with aluminum sulfate. Subsequently, various bulking promoters were added in an amount of 3% based on the pulp. Each resultant mixture was formed into a sheet with a rectangular TAPPI paper machine using an 80-mesh wire. The sheet obtained was pressed with a press at 3.5 kg/cm² for 2 minutes and dried with a drum dryer at 105°C for 1 minute. After each dried sheet was held under the conditions of 20°C and a humidity of 65% for 1 day to regulate its moisture content, it was evaluated for bulk density as a measure of paper bulkiness and for tearing strength as a measure of paper strength performance. Ten found values were averaged.

(Evaluation item and method)

Bulkiness (bulk density)

50

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[0035] The basis weight  $(g/m^2)$  and thickness (mm) of each sheet having a regulated moisture content were measured, and its bulk density  $(g/cm^3)$  was determined as a calculated value.

Equation for calculation:

[0036]

Bulkiness (bulk density) = (basis weight)/(thickness) × 0.001

• [0037] The smaller the absolute value of bulk density, the higher the bulkiness. A difference of 0.02 in bulk density is sufficiently recognized as a significant difference.

· Paper strength (tearing strength)

[0038] Each sheet having a regulated moisture content was examined according to JIS P 8116 (Testing Method for Tearing Strength of Paper and Paperboard).

Equation for calculation:

[0039]

15

Tearing strength =  $A/S \times 16$ 

Tearing strength: (gf)

A: Reading

S: Number of torn sheets

[0040] The larger the absolute value of tearing strength, the higher the paper strength. A difference of 20 gf in tearing strength is sufficiently recognized as a significant difference.

Table 1

	Deinked pulp		LBKP	
	Bulk density (g/cm <sup>3</sup> )	Tearing strength (gf)	Bulk density (g/cm <sup>3</sup> )	Tearing strength (gf)
Example 1	0.328	403	0.377	490
Example 2	0.329	435	0.376	500
Example 3	0.326	425	0.379	490
Example 4	0.334	420	0.383	485
Example 5	0.341	430	0.386	480
Example 6	0.344	410	0.384	490
Example 7	0.336	420	0.382	500
Example 8	0.348	410	0.387	485
Example 9	0.330	430	0.378	495
Example 10	0.328	440	0.379	505
Example 11	0.309	415	0.360	475
Example 12	0.307	410	0.357	470
Example 13	0.311	410	0.362	470
Example 14	0.323	415	0.371	480
Example 15	0.347	420	0.388	485
Example 16	0.307	410	0.360	465
Example 17	0.308	410	0.361	470 '
Example 18	0.314	415	0.363	475
Example 19	0.320	420	0.371	470
Example 20	0.322	425	0.373	475
Comparative Example 1	0.370	450	0.408	500

## Table 1 (continued)

	Deinked pulp		LBKP	
<del></del>	Bulk density (g/cm <sup>3</sup> )	Tearing strength (gf)	Bulk density (g/cm3)	Tearing strength (gf)
Comparative Example 2	0.372	430	0.414	490
Comparative Example 3	0.368	435	0.409	480
Comparative Example 4	0.367	425	0.410	505
Comparative Example 5	0.375	430	0.414	490
Comparative Example 6	0.330	280	0.379	345

## (Bulking promoters used)

[0041]

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	•	
	• Example 1:	Decyl alcohol/EO adduct; EOp (average number of moles of EO added; the same applies hereinafter) = 1.5.
25	• Example 2:	EO/PO block adduct of lauryl alcohol; EOp = 2.0, POp (average number of moles of PO added; the same applies hereinafter) = 0.8.
	• Example 3:	EO adduct of an octyl alcohol/decyl alcohol/lauryl alcohol/myristyl alcohol mixture (weight ratio: 8/38/30/24); EOp = 1.2.
30	• Example 4:	EO/PO random adduct of a decyl alcohol/lauryl alcohol mixture (weight ratio: 60/40); EOp=1.6, POp =0.4.
••	• Example 5:	Dobanol/EO adduct; EOp =1.0.
	• Example 6:	EO/PO random adduct of lauryl alcohol; EOp = 10.0, POp = 7.5.
	• Example 7:	EO adduct of an octyl alcohol/decyl alcohol/oleyl alcohol mixture (weight ratio: 10/60/30);
	Litample 7.	EOp =1.0.
35	• Example 8:	Nonylphenol/EO adduct; EOp =4.8.
••	• Example 9:	Mixture of decyl alcohol, polyoxyethylene (EOp = 20.0) sorbitan trioleate, and polyoxyethyl-
	Example 5.	ene hardened castor oil (EOp = 30.0) (weight ratio: 80/14/6).
	• Example 10:	Mixture of (A) EO adduct (EOp = 1.2) of a decyl alcohol/lauryl alcohol/myristyl alcohol mix-
	Example 10.	ture (weight ratio:40/30/30), (B) polyoxyethylene (EOp = 30.0) sorbitan monooleate, and (C)
40		polyoxyethylene (EOp = 25.0) hardened castor oil (weight ratio: 78/14/8).
	• Example 11:	Lauryl alcohol.
	• Example 12:	PO adduct of a lauryl alcohol/myristyl alcohol mixture (weight ratio: 50/50); POp = 5.
	• Example 13:	Mixture of lauryl alcohol, polyoxyethylene (EOp = 14) sorbitan monooleate, and polyoxyeth-
		ylene (EOp = 25) hardened castor oil (weight ratio: 80/14/6).
45	• Example 14:	EO/PO random adduct of a lauryl alcohol/myristyl alcohol mixture (weight ratio: 70/30);
	•	EOp = 2, POp = 5.
	• Example 15:	Stearyl alcohol.
	• Example 16:	Stearyl alcohol/PO adduct; POp = 10.
	Example 17:	Stearyl alcohol/PO adduct; POp = 40.
50	<ul><li>Example 18:</li></ul>	Oleyl alcohol.
	<ul> <li>Example 19:</li> </ul>	Mixture of lauryl alcohol, myristyl alcohol, and polyoxyethylene (EOp = 12) sorbitan
		monooleate (weight ratio: 68/16/16).
	• Example 20:	Mixture of lauryl alcohol and polyoxyethylene (EOp = 25) hardened castor oil (weight ratio:
		80/20).
55	<ul> <li>Comparative Example 1:</li> </ul>	
	<ul> <li>Comparative Example 2:</li> </ul>	
	O	D1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1

• Comparative Example 3: Decyl alcohol/EO adduct; EOp = 30.

• Comparative Example 4: EO/PO random adduct of lauryl alcohol; EOp = 30, POp = 30.

- Comparative Example 5: Blank (no bulking promoters).
- Comparative Example 6: Commercial bulking promoter "Bayvolume P Liquid" (fatty acid polyamide polyamine type; manufactured by Bayer AG).

#### 5 Claims

1. A paper bulking promoter which contains a compound represented by the following formula (1):

 $RO(EO)_{m}(PO)_{n}H \tag{1}$ 

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wherein R represents a linear or branched alkyl or alkenyl group having 6 to 22 carbon atoms or an alkylaryl group in which the alkyl group has 4 to 20 carbon atoms; E represents an ethylene group; P represents a propylene group; and m and n respectively indicate the average number of moles added in the ranges of  $0 \le m \le 20$  and  $0 \le n \le 50$ ; provided that  $(EO)_m(PO)_n$  may have any of block and random arrangements and may begin with any of EO and PO.

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- The paper bulking promoter as set forth in claim 1, wherein R in the formula is a linear or branched alkyl or alkenyl group having 8 to 18 carbon atoms.
- 3. The paper bulking promoter as set forth in claim 1, wherein the compound (1) is one obtained by causing a mixture of two or more ROH's to add at least either of EO and PO.
  - 4. The paper bulking promoter composition which contains the compound (1) as described in claim 1 and at least one nonionic surfactant based on a polyhydric alcohol.
- 5. The composition as set forth in claim 4, wherein the nonionic surfactant based on a polyhydric alcohol is at least one member selected among sugar alcohol/EO (ethylene oxide, the same applies hereinafter) adducts, fatty acid esters of the adducts, fatty acid esters of sugar alcohols, sugar/EO adducts, fatty acid esters of the adducts, sugar/fatty acid esters, and fat/EO adducts.
- 30 6. The composition as set forth in claim 4, wherein the nonionic surfactant based on a polyhydric alcohol is at least one member selected among fatty acid esters of sugar alcohol/EO adducts and among fat/EO adducts.
  - 7. The composition as set forth in claim 4, wherein the proportion of the compound (1) to the nonionic surfactant based on a polyhydric alcohol is from 5/5 to 10/0 (by weight).

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- 8. The composition as set forth in claim 4, wherein the proportion of the compound (1) to the nonionic surfactant based on a polyhydric alcohol is from 5/5 to 99/1 (by weight).
- 9. The composition as set forth in claim 4, wherein 0≤m≤2 and the proportion of the compound (1) to the nonionic surfactant based on a polyhydric alcohol is from 5/5 to 99/1 (by weight).
  - 10. The composition as set forth in claim 4, wherein m and n are zero and the proportion of the compound (1) to the nonionic surfactant based on a polyhydric alcohol is from 5/5 to 99/1 (by weight).
- 45 11. A process for producing a highly bulky pulp sheet which comprises adding the bulking promoter as in any one of claims 1 or 4 in any of the papermaking process steps to thereby produce a highly bulky pulp sheet which has a bulk density lower by at least 5% than the product not containing the paper bulking promoter and a tearing strength as measured according to JIS P 8116 of at least 90% of that of the product.
- 12. The process as set forth in claim 11 which comprises mixing the bulking promoter as set forth in claim 1 or 4 with water and a pulp feedstock to obtain a homogeneous slurry and subjecting the slurry to sheet forming.
  - 13. The process as set forth in claim 11 which comprises mixing an aqueous emulsion or mixture containing 10 to 100 wt.% of the bulking promoter as set forth in claim 1 or 4 with a pulp feedstock to obtain a homogeneous slurry and subjecting the slurry to sheet forming.
  - 14. A highly bulky pulp sheet obtained by the process as set forth in claim 11.

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	15. A highly bulky pulp sheet which comprises the bulking promoter as set forth in claim 1 or 4 and a pulp paper and which has a bulk density lower by at least 5% than the product not containing the paper bulking promoter and a tearing strength as measured according to JIS P 8116 of at least 90% of that of the product.		

## INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP97/02484

A. CLASSIFICATION OF SUBJECT MATTER			
Int. Cl <sup>6</sup> D21H17/06, 17/53			
	to International Patent Classification (IPC) or to both	h national classification and IPC	
	DS SEARCHED		
Micimum d	ocumentation searched (classification system followed b	ey classification symbols)	
Inc.	C16 D21H11/00-27/42		
Documentat	ion searched other than minimum documentation to the	extent that such documents are included in the	re Gelds searched
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Toro	ku Jitsuyo Shinan Koho	1994 - 1997	
Electronic di	ta base consulted during the international search (name	of data base and, where practicable, search (	terms used)
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C. DOCU	MENTS CONSIDERED TO BE RELEVANT		
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"L" docume	" document which may throw doubts on priority claim(s) or which is considered novel of cannot be considered to involve an inventive		
cited to special r	establish the publication date of another citation or other eason (as specified)	"Y" document of particular relevance; the	P.
Decoration reterring to an one disclosure, use, exhibition or other combined with one or more other such documents, such combination			
P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family			
Date of the actual completion of the international search  Date of mailing of the international search report			
September 25, 1997 (25. 09. 97) October 7, 1997 (07. 10. 97)			
Name and mailing address of the ISA/ Authorized officer			
The state of the s			
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acsimile No. Telephone No.			

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